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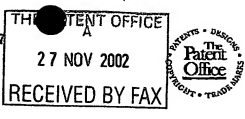
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02 E766704-1 D02884-P01/7700 0.00-0227644.2

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Your reference

P32800-/CPA/MCM

2. Patent application number (The Patent Office will fill in this part)

0227644.2

27 NOV 2002

 Full name, address and postcode of the or of each applicant (underline all surnames)

Cancer Research Technology Limited 61 Lincoln's Inn Fields London WC2A 3PX

Patents ADP number (if you know it)

United Kingdom

If the applicant is a corporate body, give the country/state of its incorporation

8497927002

UK

4. Title of the invention

"Specific Binding Members and Uses Thereof"

5. Name of your agent (If you have one)

"Address for service" in the United Kingdom to which all correspondence should be sent (including the postcode) Murgitroyd & Company

Scotland House 165-169 Scotland Street Glasgow

G5 8PL

Patents ADP number (if you know it)

1198015

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DUPLICATE

1

1 Specific Binding Members and Uses Thereof" 2 The present invention relates to specific binding 3 4 members and their use in therapy. In particular, the invention relates to specific binding members which 5 bind to CD55, their use in the modulation of 7 complement activation and the treatment of disease, . 8 for example, neoplastic disease. 9 The human complement system consists of a highly 10 11 efficient recognition and effector mechanism that consists of 30 serum or cellular components 12 13 including activated proteins, receptors and positive 14 and negative regulators. In brief, the complement 15 cascade consists of a triggering step, an 16 amplification step with a feedback loop and finally, 17 a membrane attack or lytic step. The central 18 component of the complement system is C3. Generation 19 of C3b by the classical or alternative pathway is 20 crucial for opsonisation and lysis. The classical 21 pathway is initiated when component Cl via its Clq 22 subcomponent attaches to an antibody to form an

- 1 immune complex. For the alternative pathway,
- 2 however, there is no initiating factor equivalent to
- antibody. Rather it is in a state of continuous,
- 4 low level activation as a result of spontaneous
- 5 hydrolysis of a thioester group in native C3. This
- 6 results in binding of C3 to non-specific acceptor
- 7 molecules in plasma or on cell surfaces. This can
- 8 result in the formation of C3 convertases and
- 9 creation of a feedback loop. Because of its potent
- 10 pro-flammatory and destructive capabilities, there
- is a regulatory system designed to prevent
- 12 complement activation both in the fluid phase and on
- 13 bystander tissues.

14

- 15 There are four membrane bound complement regulatory
- 16 proteins namely complement receptor 1 (CR1), CD55,
- 17 CD46 and CD59 (Liszewski et al 1996. Adv Immunol
- 18 61:201-283). Regulation is either accomplished by:

19

- 20 1. Spontaneous decay of activated proteins and
- 21 enzyme complex (i.e. short half life)
- 22 2. Destabilisation and inhibition of activation
- 23 complexes
- 24 3. Proteolytic cleavage of "activated" components.

25

- 26 CD46, CD55 and CD59 are widely expressed on many
- 27 tissues, including surface epithelia and tumour
- 28 tissues. In contrast, CR1 expression is limited to
- 29 peripheral blood cells and is therefore not directly
- 30 involved in protection of solid tumours.

1	Most tumours are of epithelial origin and, although
2	most surface epithelia express complement regulatory
3	proteins, tumours show variable expression of CD55,
4	CD46 and CD59. The majority of colorectal and
5	thyroid cancers express high levels of all three
6	complement regulatory proteins (Niehans et al., 1996
7	Am J Pathol 149:129-142; Li et al., 2001 Br. J.
8	Cancer 84:80-86; Thorsteinsson, 1998 APMIS 106:869-
9	878; Yamakawa et al., 1994 Cancer 73:2808-2817).
10	Ductal carcinoma of the breast shows the most
1.1	variation in phenotype with some tumours expressing
12	only one inhibitor while others express different
13	combinations of two or three inhibitors (Niehans et
14	al., 1996 supra; Thorsteinsson et al., 1998 supra).
15	Renal cell carcinoma has weak to moderate expression
16	of one to three inhibitors, generally CD55 and CD59
17	(Niehans et al., 1996 supra) whereas non-small cell
18	lung carcinomas and ovarian and cervical cancers
19	usually express CD59 and CD46 with variable CD55
20	immunoreactivity (Niehans et al., 1996 supra; Bjorge
21	et al., 1977 Cancer Immunol Immunother 42:185-192;
22	Simpson et al., 1997 Am J Pathol 151:1455-1467).
23	Similar results have been obtained with established
24	cell lines (Bjorge et al., 1996 supra; Gorter et al
25	1986 Lab Invest 74 1; Juhl et al., 1997 J. Surgical
26	Oncol. 64:222-230; Li et al., 2001 supra).
27	
28	All three complement regulatory proteins are
29	expressed on vascular endothelium. Their specific
30	roles during inflammation when the risk of
31	complement mediate injury may be increased remains

to be determined. CD55, but not CD46 or CD59, is

1	up-regulated on endotherial cerrs by the pro-
2	inflammatory mediators TNF α , IL-1 β , and IFN- γ , and
3	also by the MAC (membrane attack complex) and
4	thrombin. These results suggest that CD55 is of
5	critical importance in protecting endothelial cells
6	from complement during inflammation and coagulation.
7	Furthermore it has recently been shown that
8	retraction of endothelial cells exposing sub-
9	endothelial extracellular matrix is a potent inducer
10	of the alternative complement pathway releasing
11	anaphylatoxins that stimulate inflammation. As
12	tumours frequently have disregulated endothelium,
13	with exposed vessel walls, the tumour environment
14	may induce complement activation. This may be one
15	of the reasons that tumour cells over-express
16	complement regulatory receptors. However, it has
17	been shown that both tumour cells and endothelial
18	cells can actually secrete CD55 but not CD46 into
19	their extracellular matrix (ECM) (Hindmarsh and
20	Marks, 1998 J. Immunol. 160:6128-6136). Hindmarsh
21	and Marks showed that tumour but not endothelial
22	derived CD55 is functionally active and can prevent
23	deposition of C3b. However, deposition of matrix
24	CD55 could not be up-regulated by inflammatory
25	cytokines. More recently the present inventors have
26	shown that both CD55 and CD59 can be deposited into
27	extracellular matrix by both tumours and endothelial
28	cells and the latter can be considerably up-
29	regulated by the potent angiogenesis growth factor
30	VEGF (Li et al., 2001 supra). Furthermore, CD55
31	deposited by endothelial cells stimulated with VEGF
32	was shown to be functionally active. VEGF is

1 unusual, as it is the only cytokine identified to date that up-regulates both cell surface expression 2 3 and deposition of CD55 into the ECM. 4 5 As most tumours secrete high levels of VEGF to 6 induce angiogenesis they will stimulate expression 7 of CD55 on endothelial cells and within ECM. 8 Interestingly immunohistochemistry of colorectal 9 tumours with anti-CD55 monoclonal antibodies shows 10 intense staining of tumour stroma (Li et al., 2001 11 supra; Simpson et al., 1997 supra; Niehans et al., 12 1996 supra) and blood vessels (Niehans et al., 1996 13 supra). CD55 deposited within ECM is covalently 14 bound as it cannot be released by strong acids or alkalis. 15 16 17 CD55 binds C3 convertases from both the classical 18. and alternative complement pathways displacing C2b 19 and C3b respectively. It can, therefore, prevent 20 C3b deposition and inhibit the downstream assembly 21 of the membrane attack complex. CD55 has an extracellular domain that is composed of 4 22 23 contiguous short consensus (SCR) domains and a 24 threonine/serine rich region proximal to the cell 25 It has a single N-glycosylation site 26 between the first and second SCR domains and is 27 heavily O-glycosylated in the threonine and serine 28 rich regions. It is attached to the cell membrane 29 by a glycophosphoinositol (GPI) anchor and is expressed by all cells exposed to complement, 30 31 namely, red blood cells, leukocytes, endothelial and

epithelial cells. CD55 has also been detected in

1 low amounts in plasma, saliva and urine. 2 biological significance of this soluble form remains unclear as it has never been shown to be 3 functionally active. Recently it has been shown 4 5 that HeLa cells and HUVEC incorporate CD55 into 6 their extracellular matrix and that this covalently 7 linked CD55 can inhibit C3b deposition and the 8 release of the pro-inflammatory anaphylatoxin C3a 9 (Hindmarsh and Marks, 1998 supra). 10 As well as making tumour cells susceptible to in 11 12 situ complement activation, antibodies inhibiting 13 the functions of complement regulatory proteins may 14 also make tumour cells susceptible to monoclonal antibody mediated complement dependent cellular 15 16 cytotoxicity. A chimeric anti-LewisY monoclonal antibody (cH18A) mediated modest complement mediated 17 18 cell lysis of two lung adenocarcinomas cell lines. 19 However addition of antibodies that block the 20 function of CD46, CD55 and CD59 considerably enhance 21 complement mediated lysis. Use of multiple blocking 22 antibodies to the complement regulatory proteins produced more enhancement of cH18A mediated lysis 23 24 than any single antibody (Azuma et al., 1995. Scand 25 J Immunol 42:202-208). Several groups have generated bispecific antibodies with one arm targeting a 26 27 tumour cell surface antigen and the other targeting 28 the functional domain of a complement regulatory 29 protein. A bispecific antibody targeting HLA and 30 SCR3 of CD55 resulted in a 92% enhancement of C3b 31 deposition on renal tumours. Similarly in the same 32 study a bispecific antibody targeting a renal tumour

antigen and the SCR3 of CD55 resulted in a 25-400%

2 increase in C3b deposition on renal tumours and

3 rendered the cells susceptible to complement

4 mediated lysis (Blok et al., 1998 J Immunol

5 160:3437-3443). Finally when a chimeric anti-CD37

6 monoclonal antibody was used to activate the

7 classical complement pathway, a bispecific Fab'gamma

8 construct targeting a lymphoma specific antigen and

9 the CD59 functional domain increased cell lysis by

10 3-5 fold (Harris et al., 1997 Clin. Exp. Immunol.

11 107:364-371).

12

13 However, although previous studies have shown that

14 monoclonal antibodies recognising SCR3 of CD55 could

partially neutralise CD55 leading to enhanced C3b

deposition and assembly of the MAC complex, each of

these antibodies merely compete for binding to SCR3

18 with the C3 convertases and therefore only partially

19 neutralise CD55. Molecular constructs of CD55 have

20 shown that SCR3 is the active domain of CD55 and

21 that SCR2 and SCR4 are necessary to provide the

22 correct conformation for C3 binding. No role for

23 SCR1 in complement decay has been shown. However,

24 although SCR2 is necessary to provide the correct

conformation for C3 binding, studies with monoclonal

26 antibodies to single SCR domains of CD55 have shown

27 that only monoclonal antibodies that bind to SCR3

28 and not antibodies that bind to either SCR1 or SCR2

29 can neutralise CD55 (Coyne et al, 1992 J Immunol

30 149, 2906).

1 Imaging studies with the monoclonal antibody 791T/36 2 (Embleton et al 1981 Br.J. Cancer 43:582-587) in 3 osteosarcomas, ovarian and colorectal tumours 4 successfully imaged lesions as small as 1cm3 5 (Farrands et al 1982 Lancet 2:397-400; Farrands et 6 al 1983. J. of Bone and Joint Surg. 65:638-640; Armitage et al., 1985. Nucl Med Commun 6:623-631). 7 8 Furthermore autoradiography of the resected tumours showed both cell surface and intense stromal 9 10 localisation of the antibody (Armitage et al., 1984 11 Br J Surg 71:407-412). These studies illustrate that 12 an anti-CD55 antibody can effectively localise in tumours without showing any normal tissue toxicity. 13 14 In particular no detectable binding of radiolabeled 15 antibody to blood cells and only background levels of radiolabel were seen on endothelium or normal 16 tissues. The antigen recognised by 791T/36 was 17 recently identified as CD55 (Spendlove et al Eur J 18 19 Immunol. 30:2944-2953; Spendlove et al Cancer Res. 20 59:2282-2286). Using CD55/CD46 chimeric constructs 21 it was possible to map the binding site of 791T/36 22 to the first two SCR domains of CD55 with peptide 23 analysis showing that 791T/36 can bind to three distinct regions of SCR1-2 of CD55. One region is in 24 SCR1 and two are in SCR2. 25 26 27 W000/5204 discloses a method for making antibodies, 28 for example antibodies directed against decay 29 accelerating factor (DAF, using a naïve antibody phage library. Although the document refers to the 30 31 use of such antibodies in cancer diagnosis or 32 therapy, no examples are provided other than a

1	speculative example, in which antibody LU30 is
2	suggested for use in assessing overexpression of DAR
3	and for treatment of lung cancer particularly when
4	combined with cytotoxic agents.
5	
6	WO/04415 describes the production of the anti-
7	idiotype antibody 105AD7 which was raised against
8	antibody 791T/36 and speculates on potential
9	therapeutic uses of the 105AD7 antibody.
10	
11	However, to date, no therapeutically useful anti-
12	CD55 antibodies other than anti SCR3 antibodies have
13	been demonstrated. Therapeutic studies with
14	antibodies directed to other SCRs of this molecule
15	have been limited to immunoconjugated molecules.
16	(See for example US 4916213 (Koma Corporation), US
17	4925922 (Xoma Corporation) and Byers et al. 1987
18	Cancer Res 47:5042-5046). For example, Byers et al
19	describes studies with 791T/36 linked to ricin A
20	chain, showed significantly inhibition of tumour
21	growth in athymic mice. 791T/36-RTA was therefore
22	screened in a phase I clinical trial in advanced
23	colorectal cancer patients (Byers et al 1989. Cancer
24	Research 49:6153-6160). However the trial was
25	unsuccessful due to dose limiting toxicity.
26	
27	Surprisingly, the present inventors have now
28	demonstrated that, although previous studies have
29	demonstrated that antibodies which target either SCR
30	1 or SCR 2 of CD55 failed to have any neutralisation
31	effect on CD55 an antibody which targets both SCP 1

31

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and SCR2 not only effectively neutralises CD55 but 1 2 is superior to a SCR3 neutralising antibody. 3 4 Accordingly, in a first aspect, the present 5 invention provides a method of neutralisation of 6 CD55, comprising administration of a naked binding 7 member which specifically binds to SCR1 and SCR2 of 8 CD55. 9 10 By neutralising CD55, enhanced complement deposition 11 may be facilitated. Accordingly, in a second aspect, 12 the invention provides a method of enhancing 13 complement deposition on a tissue comprising 14 administration of a naked binding member which 15 specifically binds to SCR1 and SCR2 of CD55. 16 17 The methods of the invention may be used in vitro or in vivo. 18 19 20 As described above, CD55 is commonly found on many tumour cell surfaces, where it serves to inhibit 21 22 complement deposition. By neutralising such 23 molecules on tumour cells, the methods of the invention enable complement mediated attack of 24 25 tumour cells. Accordingly, in a further aspect of 26 the present invention, there is provided a method of 27 treating cancer comprising administration of a 28 therapeutically effective amount of a naked binding 29 member which specifically binds to SCR1 and SCR2 of

CD55 to a mammal in need thereof.

1	In a further aspect, there is provided the use of
2	(i) a naked binding member which binds to both SCR1
3	and SCR2 of CD55 or (ii) a nucleic acid encoding
4	said binding member in the preparation of a
5	medicament for the neutralisation of CD55.
6	
7	In a further aspect, there is provided a naked
8	binding member which binds to both SCR1 and SCR2 for
9	use in the treatment of cancer.
10	
11	In a further aspect, there is provided the use of
12	(i) a naked binding member which binds to both SCR1
13	and SCR2 of CD55 or (ii) a nucleic acid encoding
14	said binding member in the preparation of a
15	medicament for treating cancer.
16	
17	The present invention also provides a pharmaceutical
18	composition for the treatment of cancer, wherein the
19	composition comprises a naked binding member that
20	binds to both SCR1 and SCR2 of CD55.
21	
22	Specific Binding Member
23	
24	As used herein, a "binding member" is a member of a
25	pair of molecules which have binding specificity for
26	one another. The binding member is, therefore, a
27	specific binding member. The members of a binding
28	pair may be naturally derived or wholly or partially
29	synthetically produced. One member of the pair of
30	molecules may have an area on its surface, which may
31	be a protrusion or a cavity, which specifically

binds to and is therefore complementary to a

12

1 particular spatial and polar organisation of the 2 other member of the pair of molecules. Thus, the members of the pair have the property of binding 3 4 specifically to each other. Examples of types of binding pairs are antigen-antibody, biotin-avidin, 5 hormone-hormone receptor, receptor-ligand, enzyme-6 7 substrate. The present invention is concerned with 8 antigen-antibody type reactions, although a binding member of the invention and for use in the invention 9 10 may be any moiety which can bind to both SCR1 and SCR2 of CD55. 11 12 13 As used herein, "naked" means that the binding 14 member of or for use in the present invention is not bound to, for example conjugated with, any agent, 15 16 for example ricin, having anti-tumour properties. 17 18 Antibodies 19 20 An "antibody" is an immunoglobulin, whether natural or partly or wholly synthetically produced. 21 term also covers any polypeptide, protein or peptide 22 having a binding domain which is, or is homologous 23 24 to, an antibody binding domain. These can be derived from natural sources, or they may be partly 25 26 or wholly synthetically produced. Examples of 27 antibodies are the immunoglobulin isotypes and their isotypic subclasses and fragments which comprise an 28 29 antigen binding domain such as Fab, scFv, Fv, dAb, 30 Fd; and diabodies.

- 1 The binding member of the invention may be an
- 2 antibody such as a monoclonal or polyclonal
- antibody, or a fragment thereof. The constant region
- 4 of the antibody may be of any class including, but
- 5 not limited to, human classes IgG, IgA, IgM, IgD and
- 6 IgE. The antibody may belong to any sub class e.g.
- 7 IgG1, IgG2, IgG3 and IgG4. IgG1 is preferred. In
- 8 preferred embodiments the antibody is 791T/36
- 9 produced by the cell line deposited with ATCC under
- 10 accession no. HB9173.

- 12 As antibodies can be modified in a number of ways,
- 13 the term "antibody" should be construed as covering
- 14 any binding member or substance having a binding
- 15 domain with the required specificity. Thus, this
- 16 term covers antibody fragments, derivatives,
- 17 functional equivalents and homologues of antibodies,
- 18 including any polypeptide comprising an
- immunoglobulin binding domain, whether natural or
- 20 wholly or partially synthetic. Chimeric molecules
- 21 comprising an immunoglobulin binding domain, or
- 22 equivalent, fused to another polypeptide are
- 23 therefore included. Cloning and expression of
- 24 chimeric antibodies are described in EP-A-0120694
- 25 and EP-A-0125023.

- 27 It has been shown that fragments of a whole antibody
- 28 can perform the function of binding antigens.
- 29 Examples of such binding fragments are (i) the Fab
- 30 fragment consisting of VL, VH, CL and CH1 domains;
- 31 (ii) the Fd fragment consisting of the VH and CH1
- 32 domains; (iii) the Fv fragment consisting of the VL

1 and VH domains of a single antibody; (iv) the dAb fragment (Ward, E.S. et al., Nature 341:544-546 2 3 (1989)) which consists of a VH domain; (v) isolated 4 CDR regions; (vi) F(ab')2 fragments, a bivalent 5 fragment comprising two linked Fab fragments (vii) 6 single chain Fv molecules (scFv), wherein a VH 7 domain and a VL domain are linked by a peptide 8 linker which allows the two domains to associate to form an antigen binding site (Bird et al., Science 9 10 242:423-426 (1988); Huston et al., PNAS USA 85:5879-5883 (1988)); (viii) bispecific single chain Fv 11 dimers (PCT/US92/09965) and (ix) "diabodies", 12 multivalent or multispecific fragments constructed 13 14 by gene fusion (WO94/13804; P. Hollinger et al., Proc. Natl. Acad. Sci. USA 90:6444-6448 (1993)). 15 16 17 A fragment of an antibody or of a polypeptide for 18 use in the present invention, for example, a fragment of the 791T/36 antibody, generally means a 19 20 stretch of amino acid residues of at least 5 to 7 contiguous amino acids, often at least about 7 to 9 21 22 contiguous amino acids, typically at least about 9 23 to 13 contiguous amino acids, more preferably at 24 least about 20 to 30 or more contiguous amino acids and most preferably at least about 30 to 40 or more 25 26 consecutive amino acids. A preferred group of 27 fragments are those which include all or part of the 28 CDR regions of monoclonal antibody 791T/36. A 29 preferred group of fragments are those which include 30 all or part of the CDR regions of monoclonal 31 antibody 791T/36.

1	A "derivative" of such an antibody or polypeptide,
2	or of a fragment of a 791T/36 antibody means an
3	antibody or polypeptide modified by varying the
4	amino acid sequence of the protein, e.g. by
5	manipulation of the nucleic acid encoding the
6	protein or by altering the protein itself. Such
7	derivatives of the natural amino acid sequence may
8	involve insertion, addition, deletion and/or
9	substitution of one or more amino acids, preferably
10	while providing a peptide having anti-CD55 activity,
11	for example, CD55 neutralisation activity.
12	Preferably such derivatives involve the insertion,
13·	addition, deletion and/or substitution of 25 or
14	fewer amino acids, more preferably of 15 or fewer,
15	even more preferably of 10 or fewer, more preferably
16	still of 4 or fewer and most preferably of 1 or 2
17	amino acids only.
18	· ·
19	The term "antibody" includes antibodies which have
20	been "humanised". Methods for making humanised
21	antibodies are known in the art. Methods are
22	described, for example, in Winter, U.S. Patent No.
23	5,225,539. A humanised antibody may be a modified
24	antibody having the hypervariable region of a
25	monoclonal antibody such as 791T/36 and the constant
26	region of a human antibody. Thus the binding member
27	may comprise a human constant region.
28	
29	The variable region other than the hypervariable
30	region may also be derived from the variable region
31	of a human antibody and/or may also be derived from
32	a monoclonal antibody such as 791T/36 In such

1 case, the entire variable region may be derived from 2 murine monoclonal antibody 791T/36 and the antibody 3 is said to be chimerised. Methods for making 4 chimerised antibodies are known in the art. 5 methods include, for example, those described in 6 U.S. patents by Boss (Celltech) and by Cabilly 7 (Genentech). See U.S. Patent Nos. 4,816,397 and 8 4,816,567, respectively. 9 10 It is possible to take monoclonal and other antibodies and use techniques of recombinant DNA 11 12 technology to produce other antibodies or chimeric 13 molecules which retain the specificity of the 14 original antibody. Such techniques may involve introducing DNA encoding the immunoglobulin variable 15 16 region, or the complementary determining regions 17 (CDRs), of an antibody to the constant regions, or 18 constant regions plus framework regions, of a 19 different immunoglobulin. See, for instance, EP-A-184187, GB 2188638A or EP-A-239400. A hybridoma or 20 other cell producing an antibody may be subject to 21 genetic mutation or other changes, which may or may 22 23 not alter the binding specificity of antibodies 24 produced. 25 In preferred embodiments of the invention, the 26 binding member binds to CD55 SCR1 (amino acids 83-27 28 93) and SCR2 (amino acids 101-112 and amino acids 29 145-157) of the sequences shown in Figure 1b. 30 31 The binding member may comprise one or more of the CDRs of the antibody, or a fragment thereof, 32

1 produced by the cell line deposited at ATCC under accession number HB9173. 2

3

4

As described above, in a preferred embodiment of the invention, the binding member is the antibody 5 791T/36 produced by the hybridoma cell deposited 6 7 under ATCC accession number HB9173. As used herein, reference to "791T/36" includes sequences which show 8 substantial homology with 791T/36. Preferably the 9 degree of homology between 791T/36 complementary 10 11 determining regions (CDRs) and the CDRs of other antibodies will be at least 60%, more preferably 12

70%, further preferably 80%, even more preferably

90% or most preferably 95%.

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The percent identity of two amino acid sequences or of two nucleic acid sequences may be determined by aligning the sequences for optimal comparison purposes (e.g., gaps can be introduced in the first sequence for best alignment with the sequence) and comparing the amino acid residues or nucleotides at corresponding positions. The "best alignment" is an alignment of two sequences which results in the highest percent identity. The percent identity is determined by the number of identical amino acid residues or nucleotides in the sequences being compared (i.e., % identity = number of identical positions/total number of positions x 100).

28 29 30

31

32

The determination of percent identity between two sequences can be accomplished using a mathematical algorithm known to those of skill in the art.

- example of a mathematical algorithm for comparing
- 2 two sequences is the algorithm of Karlin and
- 3 Altschul (1990) Proc. Natl. Acad. Sci. USA
- 4 87:2264-2268, modified as in Karlin and Altschul
- 5 (1993) Proc. Natl. Acad. Sci. USA 90:5873-5877. The
- 6 NBLAST and XBLAST programs of Altschul, et al.
- 7 (1990) J. Mol. Biol. 215:403-410 have incorporated
- 8 such an algorithm. BLAST nucleotide searches can be
- 9 performed with the NBLAST program, score = 100,
- wordlength = 12 to obtain nucleotide sequences
- 11 homologous to nucleic acid molecules of the
- 12 invention. BLAST protein searches can be performed
- with the XBLAST program, score = 50, wordlength = 3
- 14 to obtain amino acid sequences homologous to protein
- 15 molecules of the invention. To obtain gapped
- 16 alignments for comparison purposes, Gapped BLAST can
- be utilised as described in Altschul et al. (1997)
- 18 Nucleic Acids Res. 25:3389-3402. Alternatively,
- 19 PSI-Blast can be used to perform an iterated search
- 20 which detects distant relationships between
- 21 molecules (Id.). When utilising BLAST, Gapped
- 22 BLAST, and PSI-Blast programs, the default
- 23 parameters of the respective programs (e.g., XBLAST
- 24 and NBLAST) can be used. See
- 25 http://www.ncbi.nlm.nih.gov.

- 27 Another example of a mathematical algorithm utilised
- 28 for the comparison of sequences is the algorithm of
- 29 Myers & Miller, CABIOS (1989). The ALIGN program
- 30 (version 2.0) which is part of the CGC sequence
- 31 alignment software package has incorporated such an
- 32 algorithm. Other algorithms for sequence analysis

32

1	known in the art include ADVANCE and ADAM as
2	described in Torellis & Robotti (1994) Comput. Appl
3	Biosci., 10:3-5; and FASTA described in Pearson &
4	Lipman (1988) Proc. Natl. Acad. Sci. 85:2444-8.
5	Within FASTA, ktup is a control option that sets th
6	sensitivity and speed of the search.
7	
8	Where high degrees of sequence identity are present
9	there will be relatively few differences in amino
ĿO	acid sequence. Thus for example they may be less
L1 .	than 20, less than 10, or even less than 5
12	differences.
13	
14	The present inventors have shown that antibodies
15	directed to SCR1 and SCR2 of CD55, for example
16	791T/36 antibodies and fragments and derivatives
17	thereof can be used as cancer therapeutics to
18	inactivate CD55 and make tumour cells susceptible t
19	complement mediated attack. This is exemplified by
20	localisation of the antibody within tumours of
21	cancer patients and their subsequent enhanced
22	survival (see the Examples). Accordingly the
23	invention further provides the use of naked
24	"fragments" or "derivatives" of 791T/36 or other
25	polypeptides of the "791T/36" family which bind to
26	both SCR1 and SCR2 CD55 epitopes in the preparation
27	of an agent for treating cancer.
28	
29	The binding members may be administered alone or in
30	combination with one or more further agents. Thus,

the present invention further provides products

comprising a naked binding member, which binds to

1	both SCR1 and SCR2 of CD55, and an active agent as a
2	combined preparation for simultaneous, separate or
3	sequential use in the treatment of cancer. Active
4 ·	agents may include chemotherapeutic agents
5	including, Doxorubicin, taxol, 5-Fluorouracil (5
6	FU), Leucovorin, Trinotecan, Mitomycin C,
7	Oxaliplatin, Raltitrexed, Tamoxifen and Cisplatin
8	which may operate synergistically with the binding
9	member of the present invention. Other active agents
10	may include suitable doses of pain relief drugs such
11	as non-steroidal anti-inflammatory drugs (e.g.
12	aspirin, paracetamol, ibuprofen or ketoprofen) or
13	opiates such as morphine, or anti-emetics. The
14	ability of the binding member to synergise with an
15	active agent to enhance tumour killing may not be
16	due to immune effector mechanisms but rather may be
17	a direct consequence of inactivating CD55 allowing
18	enhanced complement deposition and complement lysis.
19	The binding member of the invention may carry a
20	detectable label.
21	
22	Treatment
23	
24	"Treatment" includes any regime that can benefit a
25	human or non-human animal. The treatment may be in
26	respect of an existing condition or may be
27	prophylactic (preventative treatment). Treatment may
28	include curative, alleviation or prophylactic

29

effects.

31 "Treatment of cancer" includes treatment of

32 conditions caused by cancerous growth and includes

Т	the treatment of neoplastic growths or tumours.
2	Examples of tumours that can be treated by the
3	system of the invention are, for instance, sarcomas,
4	including osteogenic and soft tissue sarcomas,
5	carcinomas, e.g., breast-, lung-, bladder-, thyroid-
6	, prostate-, colon-, rectum-, pancreas-, stomach-,
7	liver-, uterine-, cervical and ovarian carcinoma,
8	lymphomas, including Hodgkin and non-Hodgkin
9	lymphomas, neuroblastoma, melanoma, myeloma, Wilms
10	tumor, and leukemias, including acute lymphoblastic
11	leukaemia and acute myeloblastic leukaemia, gliomas
12	and retinoblastomas.
13	•
14	The binding member may, upon binding to SCR1 and
15	SCR2 of CD55 present on cancerous cells or tissues,
16	including tumour and non-tumour cells, neutralise
17	CD55 and enhance complement deposition and
18	complement mediated lysis of these cells.
19	
20	The compositions and methods of the invention may be
21	particularly useful in the treatment of existing
22	cancer and in the prevention of the recurrence of
23	cancer after initial treatment or surgery.
24	
25	Administration
26	
27	Binding members of the present invention may be
28	administered alone but will preferably be
29	administered as a pharmaceutical composition, which
30	will generally comprise a suitable pharmaceutical
31	excipient, diluent or carrier selected dependent on
32	the intended route of administration.

Binding members of the present invention may be 1 administered to a patient in need of treatment via 2 any suitable route. The precise dose will depend 3 upon a number of factors, including the precise 4 nature of the member (e.g. whole antibody, fragment 5 or diabody), and the nature of the detectable label attached to the member. 7 8 Some suitable routes of administration include (but 9 are not limited to) oral, rectal, masal, topical 10 11 (including buccal and sublingual), vaginal or parenteral (including subcutaneous, intramuscular, 12 intravenous, intradermal, intrathecal and epidural) 13 14 administration. Intravenous administration is 15 preferred. 16 It is envisaged that injections (intravenous) will 17 be the primary route for therapeutic administration 18 19 of the compositions although delivery through a catheter or other surgical tubing is also envisaged. 20 Liquid formulations may be utilised after 21 reconstitution from powder formulations. 22 23 For intravenous, injection, or injection at the site 24 25 of affliction, the active ingredient will be in the form of a parenterally acceptable aqueous solution 26 which is pyrogen-free and has suitable pH, 27 isotonicity and stability. Those of relevant skill 28 in the art are well able to prepare suitable 29 solutions using, for example, isotonic vehicles such 30 as Sodium Chloride Injection, Ringer's Injection, 31

Lactated Ringer's Injection. Preservatives,

stabilisers, buffers, antioxidants and/or other additives may be included, as required.

3

4 Pharmaceutical compositions for oral administration may be in tablet, capsule, powder or liquid form. A 5 tablet may comprise a solid carrier such as gelatin 6 7 or an adjuvant. Liquid pharmaceutical compositions generally comprise a liquid carrier such as water, 8 9 petroleum, animal or vegetable oils, mineral oil or synthetic oil. Physiological saline solution, 10 11 dextrose or other saccharide solution or glycols such as ethylene glycol, propylene glycol or 12

polyethylene glycol may be included.

13 14

30

31 32

15 The composition may also be administered via . 16 microspheres, liposomes, other microparticulate 17 delivery systems or sustained release formulations 18 placed in certain tissues including blood. examples of sustained release carriers include 19 semipermeable polymer matrices in the form of shared 20 articles, e.g. suppositories or microcapsules. 21 22 Implantable or microcapsular sustained release 23 matrices include polylactides (US Patent No. 3, 773, 24 919; EP-A-0058481) copolymers of L-glutamic acid and 25 gamma ethyl-L-glutamate (Sidman et al, Biopolymers 26 22(1): 547-556, 1985), poly (2-hydroxyethyl-27 methacrylate) or ethylene vinyl acetate (Langer et 28 al, J. Biomed. Mater. Res. 15: 167-277, 1981, and 29 Langer, Chem. Tech. 12:98-105, 1982).

containing the polypeptides are prepared by well-known methods: DE 3,218, 121A; Epstein et al, PNAS

USA, 82: 3688-3692, 1985; Hwang et al, PNAS USA, 77:

1	4030-4034, 1980; EP-A-0052522; E-A-0036676; EP-A-
2	0088046; EP-A-0143949; EP-A-0142541; JP-A-83-11808;
3	US Patent Nos 4,485,045 and 4,544,545. Ordinarily,
4	the liposomes are of the small (about 200-800
5	Angstroms) unilamellar type in which the lipid
6	content is greater than about 30 mol. % cholesterol,
7	the selected proportion being adjusted for the
8	optimal rate of the polypeptide leakage.
9	·
10	Examples of the techniques and protocols mentioned
11	above and other techniques and protocols which may
12	be used in accordance with the invention can be
13	found in Remington's Pharmaceutical Sciences, 16th
14	edition, Oslo, A. (ed), 1980.
15	
16	The composition may be administered in a localised
17	manner to a tumour site or other desired site or may
18	be delivered in a manner in which it targets tumour
19	or other cells. Targeting therapies may be used to
20	deliver the active agent more specifically to
21	certain types of cell, by the use of targeting
22	systems such as antibody or cell specific ligands.
23	Targeting may be desirable for a variety of reasons
24	for example if the agent is unacceptably toxic, or
25	if it would otherwise require too high a dosage, or
26	if it would not otherwise be able to enter the
27	target cells.
28	
29	Pharmaceutical Compositions
30	
21	As described above, the present invention extends to

a pharmaceutical composition for the treatment of

ı	cancer, the composition comprising a naked binding
2	member which binds to both SCR1 and SCR2 of CD55.
3	Pharmaceutical compositions according to the present
4	invention, and for use in accordance with the
5	present invention may comprise, in addition to
6	active ingredient, a pharmaceutically acceptable
7	excipient, carrier, buffer stabiliser or other
8	materials well known to those skilled in the art.
9 ·	Such materials should be non-toxic and should not
.0	interfere with the efficacy of the active
11	ingredient. The precise nature of the carrier or
12	other material will depend on the route of
13	administration, which may be oral, or by injection,
14	e.g. intravenous.
15	
16	The formulation may be a liquid, for example, a
17	physiologic salt solution containing non-phosphate
18	buffer at pH 6.8-7.6, or a lyophilised powder.
19	
20	Dose
21	
22	The compositions are preferably administered to an
23	individual in a "therapeutically effective amount",
24	this being sufficient to show benefit to the
25	individual. The actual amount administered, and
26	rate and time-course of administration, will depend
27	on the nature and severity of what is being treated
28	Prescription of treatment, e.g. decisions on dosage
29	etc, is ultimately within the responsibility and at
30	the discretion of general practitioners and other
31	medical doctors, and typically takes account of the

disorder to be treated, the condition of the

1	individual patient, the site of delivery, the method
2	of administration and other factors known to
3	practitioners.
4	
5	The optimal dose can be determined by physicians
6	based on a number of parameters including, for
7	example, age, sex, weight, severity of the condition
ន	being treated, the active ingredient being
9	administered and the route of administration. In
10	general, a serum concentration of polypeptides and
11	antibodies that permits saturation of receptors is
12	desirable. A concentration in excess of
13	approximately 0.1nM is normally sufficient. For
14	example, a dose of 100mg/m2 of antibody provides a
15	serum concentration of approximately 20nM for
16	approximately eight days.
17	
18	As a rough guideline, doses of antibodies may be
19	given weekly in amounts of 10-300mg/m2. Equivalent
20	doses of antibody fragments should be used at more
21	frequent intervals in order to maintain a serum
22	level in excess of the concentration that permits
23	saturation of CD55.
24	
25	Production of Binding Members
26	·
27	The binding members of and for use in the present
28	invention may be generated wholly or partly by
29	chemical synthesis. The binding members can be
30	readily prepared according to well-established,
31	standard liquid or, preferably, solid-phase peptide
32 ·	synthesis methods, general descriptions of which are

1	broadly available (see, for example, in J.M. Stewart
2	and J.D. Young, Solid Phase Peptide Synthesis, 2nd
3	edition, Pierce Chemical Company, Rockford, Illinois
4	(1984), in M. Bodanzsky and A. Bodanzsky, The
5	Practice of Peptide Synthesis, Springer Verlag, New
6	York (1984); and Applied Biosystems 430A Users
7	Manual, ABI Inc., Foster City, California), or they
8	may be prepared in solution, by the liquid phase
9	method or by any combination of solid-phase, liquid
10	phase and solution chemistry, e.g. by first
11	completing the respective peptide portion and then,
12	if desired and appropriate, after removal of any
13	protecting groups being present, by introduction of
14	the residue X by reaction of the respective carbonic
15	or sulfonic acid or a reactive derivative thereof.
16	. ••
17	Another convenient way of producing a binding member
18	suitable for use in the present invention is to
19	express nucleic acid encoding it, by use of nucleic
20	acid in an expression system. Thus the present
21	invention further provides the use of an isolated
22	nucleic acid encoding a naked binding member which
23	binds to both SCR1 and SCR2 of CD55 in the
24	preparation of an agent for treating cancer.
25	
26	Nucleic acid for use in accordance with the present
27	invention may comprise DNA or RNA and may be wholly
28	or partially synthetic. In a preferred aspect,
29	nucleic acid for use in the invention codes for a
30	binding member of the invention as defined above.
31	The skilled person will be able to determine

substitutions, deletions and/or additions to such

nucleic acids which will still provide a binding member of the present invention.

3 Nucleic acid sequences encoding a binding member for 4 5 use with the present invention can be readily 6 prepared by the skilled person using the information 7 and references contained herein and techniques known 8 in the art (for example, see Sambrook, Fritsch and 9 Maniatis, "Molecular Cloning", A Laboratory Manual, 10 Cold Spring Harbor Laboratory Press, 1989, and 11 Ausubel et al, Short Protocols in Molecular Biology, 12 John Wiley and Sons, 1992), given the nucleic acid 13 sequences and clones available. These techniques 14 include (i) the use of the polymerase chain reaction 15 (PCR) to amplify samples of such nucleic acid, e.g. 16 from genomic sources, (ii) chemical synthesis, or 17 (iii) preparing cDNA sequences. DNA encoding 18 antibody fragments may be generated and used in any 19 suitable way known to those of skill in the art, 20 including by taking encoding DNA, identifying 21 suitable restriction enzyme recognition sites either 22 side of the portion to be expressed, and cutting out 23 said portion from the DNA. The portion may then be 24 operably linked to a suitable promoter in a standard 25 commercially available expression system. 26 recombinant approach is to amplify the relevant 27 portion of the DNA with suitable PCR primers. 28 Modifications to the sequences can be made, e.g. 29 using site directed mutagenesis, to lead to the expression of modified peptide or to take account of 30 31 codon preferences in the host cells used to express

the nucleic acid.

1	The nucleic acid may be comprised as constructs in
2	the form of a plasmid, vector, transcription or
3	expression cassette which comprises at least one
4	nucleic acid as described above. The construct may
5	be comprised within a recombinant host cell which
6	comprises one or more constructs as above.
7	Expression may conveniently be achieved by culturing
8	under appropriate conditions recombinant host cells
9	containing the nucleic acid. Following production
10	by expression a specific binding member may be
11	isolated and/or purified using any suitable
12	technique, then used as appropriate.
13	•
14	Binding members-encoding nucleic acid molecules and
15	vectors for use in accordance with the present
16	invention may be provided isolated and/or purified,
17	e.g. from their natural environment, in
18	substantially pure or homogeneous form, or, in the
19	case of nucleic acid, free or substantially free of
20	nucleic acid or genes origin other than the sequence
21	encoding a polypeptide with the required function.
22	
23	Systems for cloning and expression of a polypeptide
24	in a variety of different host cells are well known
25	Suitable host cells include bacteria, mammalian
26	cells, yeast and baculovirus systems. Mammalian
27	cell lines available in the art for expression of a

heterologous polypeptide include Chinese hamster

preferred bacterial host is E. coli.

ovary cells, HeLa cells, baby hamster kidney cells,

NSO mouse melanoma cells and many others. A common,

31 32

28

29

The expression of antibodies and antibody fragments 1 in prokaryotic cells such as E. coli is well 2 established in the art. For a review, see for 3 example Plückthun, Bio/Technology 9:545-551 (1991). 4 Expression in eukaryotic cells in culture is also 5 available to those skilled in the art as an option б for production of a binding member, see for recent 7 review, for example Reff, Curr. Opinion Biotech. 8 4:573-576 (1993); Trill et al., Curr. Opinion 9 Biotech. 6:553-560 (1995). 10 11 Suitable vectors can be chosen or constructed, 12 containing appropriate regulatory sequences, 13 including promoter sequences, terminator sequences, 14 polyadenylation sequences, enhancer sequences, 15 marker genes and other sequences as appropriate. 16 Vectors may be plasmids, viral e.g. 'phage, or 17 phagemid, as appropriate. For further details see, 18 for example, Sambrook et al., Molecular Cloning: A 19 Laboratory Manual: 2nd Edition, Cold Spring Harbor 20 Laboratory Press (1989). Many known techniques and 21 protocols for manipulation of nucleic acid, for 22 example in preparation of nucleic acid constructs, 23 mutagenesis, sequencing, introduction of DNA into 24 cells and gene expression, and analysis of proteins, 25 are described in detail in Ausubel et al. eds., 26 Short Protocols in Molecular Biology, 2nd Edition, 27 John Wiley & Sons (1992). 28 29 The nucleic acid may be introduced into a host cell 30 . by any suitable means. The introduction may employ 31 any available technique. For eukaryotic cells,

1	suitable techniques may include calcium phosphate
2	transfection, DEAE-Dextran, electroporation,
3	liposome-mediated transfection and transduction
4	using retrovirus or other virus, e.g. vaccinia or,
5	for insect cells, baculovirus. For bacterial cells,
6	suitable techniques may include calcium chloride
7	transformation, electroporation and transfection
8	using bacteriophage.
9	•
10	Marker genes such as antibiotic resistance or
11	sensitivity genes may be used in identifying clones
12	containing nucleic acid of interest, as is well
13	known in the art.
14	
15	The introduction may be followed by causing or
16	allowing expression from the nucleic acid, e.g. by
17	culturing host cells under conditions for expression
18	of the gene.
19	•
20	The nucleic acid may be integrated into the genome
21	(e.g. chromosome) of the host cell. Integration may
22	be promoted by inclusion of sequences which promote
23	recombination with the genome in accordance with
24	standard techniques. The nucleic acid may be on an
25	extra-chromosomal vector within the cell, or
26	otherwise identifiably heterologous or foreign to
27	the cell.
28	
29	Assays
30	
31	The invention further provides assays for

identification of further agents, for example

-	and poorter that tan be used for the emissionment of
2	complement deposition on a cell sample or tissue and
3	which can optionally be used in the treatment of
4 ·	cancer.
5	•
6	In a preferred aspect, the assay comprises an assay
7	method for identification of an agent capable of
8	inhibiting CD55 comprising steps:
9	·
10	a) bringing into contact a candidate agent with at.
11	least a portion of SCR1 and SCR2 of CD55; and
12	
13	b) determining binding of said candidate agent to
14	both SCR1 and SCR2.
15	
16	In a further embodiment, the assay method comprises
17	a method for identification of an agent capable of
18	inhibiting CD55 comprising:
19	
20	(a) bringing into contact a candidate agent with at
21	least a portion of SCR1 and SCR2 of CD55 in the
22	presence of a naked binding member which in the
23	absence of the candidate agent is capable of
24	binding both SCR1 and SCR2 of CD55; and
25	·
26	(b) determining the extent to which the candidate
27	agent inhibits binding of the naked binding
28	member to SCR1 and SCR2 of CD55.
29	
30	The assays may further comprise the step of
31	selecting a candidate agent which binds both SCR1
32	and SCR2 of CD55; and/or the step of determining

32

1	the amount of complement deposition on a cell sample
2	in the presence and absence of the candidate agent.
3	·
4	In preferred embodiments of the assays of the
5	invention, the portion of SCR1 and SCR2 of CD55
6	comprises amino acids 83-93, 101-112 and 145-157 of
7	the sequences shown in Figure 1b.
8	·
9	The present invention further provides a screening
10	method comprising the step of screening a library of
11	candidate agents for the ability to inhibit the
12	binding of a naked binding member to both SCR1 and
13	SCR2 of CD55.
14	
15	The assay of the invention may be a screen , whereby
 16	a number of candidate agents are tested.
17	Accordingly, any suitable technique for screening
18	compounds known to the person skilled in the art may
19	be used. The screen may be a high-throughput
20	screen. For example, WO84/03564 describes a method
21	in which large numbers of peptides are synthesised
22	on a solid substrate and reacted with an agent and
23	washed. Bound entities are detected.
24	
25	The invention also contemplates the use of
26	competitive drug screening assays in which
27	neutralising antibodies such as 791T/36 capable of
28	binding SCR1 and 2 of CD55 specifically compete with
29	a test compound for binding to SCR1 and 2 of CD55.
30	•

Agents identified by the screening method of the

present invention and their use in the manufacture

1 of a medicament for the treatment of cancer are also 2 contemplated by the invention. 3 4 Preferred features of each aspect of the invention 5 are as for each of the other aspects mutatis mutandis. 8 The invention will now be described further in the 9 following non-limiting examples. Reference is made 10 to the accompanying drawings in which: 11 12 Figure la represents the translated CDR sequences of 13 VK and VH cDNAs from 105AD7 hybridoma. Uppercase 14 letters represent the CDR regions, the lower case letters are the adjacent framework amino acids. 15 16 17 Figure 1b shows alignment of the three CDR peptides 18 with CD55. The amino acid numbering is taken from 19 the full-length sequence of CD55 including the 20 leader sequence. CD55 peptides used in subsequent assays are shown underlined. Bullets (•) represent 21 22 amino acid identity whereas amino acids with similar 23 physicochemical properties are marked as (). 24 Figure 2 illustrates a C3b complement deposition 25 assay. 791T cells were incubated with human serum as 26 a source of complement. C3b deposition was measured 27 using rabbit anti-C3b FITC labelled antibody in the 28 29 presence of blocking (216), non blocking (220) or 30 test antibody 791T/36. Fluorescence was quantified

by a FACScan flow cytomemeter and is present as mean

linear fluorescence (MLF).

1	Example 1 CD55 Neutralisation Assay
2	
3	Purified CD55 antigen was obtained by
4	immunoaffinity-matrix purification from octyl-
5 .	glucoside-solublised 791T cells. CD55 cDNA was
6	cloned and sequenced using primers based on protein
7	sequence data obtained from the purified antigen
8	(Spendlove et al., 1999 Cancer Res 59, 2282). The
9	DNA sequence obtained was identical to that
10	identified by Caras et al and present on the Genbank
11	database (Accession No. M31516).
12.	
13	Cells
14	
15	791T is an osteosarcoma cell line which was grown in
16	RPMI (Gibco, BRL, Paisley, and UK) supplemented with
17	10% heat inactivated fetal calf serum.
18 .	
19	Monoclonal Antibodies
20 21	Monoclonal antibodies 791T/36 (IgG2b anti-791Tgp72;
22	Embleton et al 1981Br.J. Cancer 43:582-587), BRIC
23	216 (IgG1 anti-SCR 3 of CD55; Tate et al 1989
24	Biochem J 261, 489), BRIC 220 (IgG1 anti-SCR 1 of
25	CD55, Tate et al 1989 Biochem J 261, 489), BRIC 110
26	(IgG1 anti-SCR 2 of CD55; Spring et al., 1987
27	Immunology 62 377; Coyne et al, 1992 J Immunol 149,
28	2906) have been reported previously. The BRIC
29	antibodies were purchased from the Blood Group
30	Reference laboratory (Bristol, UK).
31	
32	

1	Methods
2	·
3	791T tumour cells that over-express CD55 were washed
4	with media containing 10% FCS and resuspended at a
5	density of 1x 10 ⁵ cells per100µl. Primary antibody
6	was incubated with 3x sample volume (3 $ imes 10^5$
7	cells/300µl) at a concentration of 50µg/ml. Primary
8	antibodies were positive control antibody , 216
9	(anti-SCR3), negative control antibody 220 (anti-
10	SCR1) and test antibody, 791T/36 (anti-SCR1 and 2).
11	Cells and antibodies were incubated for 1 hr at 4°C
12	prior to washing in PBS. Samples were split into 3
13	samples of 100µl per tube. Human Serum was added as
14	a source of complement to total concentration of 5%
15	(Not Heat Inactivated). Tubes were inverted several
16	times and incubate at 37°C for 2 hours, mixing every
17	30 min. Cells were washed twice in PBS prior to
18	addition of polyclonal rabbit anti human C3c FITC
19	conjugated antibody (1/100) to a final volume of
20	100µl. Cells were incubated for 1 hour at 4°C prior
21	to washing twice in PBS and resuspending in 200 μ l of
22	1% cell fix.
23	
24	Results
25	
26 .	Figure 1 shows that in the presence of a non-
27	blocking antibody 220 C3b is deposited onto 791T
28	cells at modest levels (MLF 200). In the presence of
29	the CD55 neutralising antibody, 216, enhanced C3b
30	deposition is observed (MLF 350). However in the
31	presence of monoclonal antibody 791T/36 even greater

_	revers of con are deposited (mirozo). This suggests
2	that although 216 is an effective competitor with C3
3	convertase for binding to SCR3. binding of 791T/36
4	to SCR1 and SCR2 domains functionally inactivates
5	CD55 leading to a 250% increase in C3b deposition.
6	
7	Example 2. Long term survival of recurrent
8	colorectal cancer patients receiving radiolabelled
9	791T/36 for tumour imaging.
10	•
11	Antibody and Labelling
12	
13	Hybridoma 791T/36 clone 3 is the source of antibody
14	(791T/36, IgG2b isotype). Ascitic fluid from mice
15	in which the hybridoma was developing was applied to
16	a protein A-"Sepharose" column in pH 7.5 0.1 mol/1
17	citrate phosphate buffer and the column was
18	thoroughly washed. Bound immunoglobulins were
19	eluted stepwise at pH 6.0, 5.0, 4.5 and 3.0 and
20	these were then dialysed against phosphate-buffered
21	saline. The dialysate was then centrifuged at
22	1000000g for 1 h, filtered through a 0.22 µm Millex
23	"Millipore" filter, and stored at -70° C at a protein
24	concentration of lmg/ml. The preparation contained
25	only IgG2b as assessed by immunodiffusion tests with
26	mouse immunoglobulin typing antisera (Miles
27	Laboratories, Stoke Poges, Bucks.) and was pyrogen-
28	free (Boots Pharmaceuticals, Notts).
29	
30	Batches of the antibody preparation were labelled
31	with 131 by means of "lodogen" reagent. Non-bound
32	iodine was removed by gel filtration on sephadex

- 1 G25. Labelled preparations were diluted into saline
- 2 containing 1% serum albumin and sterilised by Millex
- 3 filtration.

- 5 72 patients with recurrent colorectal cancer were
- 6 imaged with the radiolabelled monoclonal antibody
- 7 791T/36. Patients received an id dose of 10µg of
- 8 antibody followed by an intravenous dose of 200µg.
- 9 2dl of preparation containing 200µg of antibody and
- 10 approximately 70MBq 131I was infused into an
- 11 antecubital vein of each patient over 30 min.

12

- 13 Survival was followed for 7 years and compared to a
- 14 contemporary group of recurrent colorectal cancer
- 15 patients. There were 12 long term survivors (16%)
- 16 in the patients who had received 791T/36 where as in
- 17 contrast only 1 out of 89 patients survived 7 years
- in the contemporary group (p> 0.001).

19

- 20 Table 1: Survival of colorectal cancer patients
- 21 receiving 791T/36 antibody.

22

Patients	Survival	Death
Imaged with 791T/36	12	60
Contemporary controls	1	88

- 24 These results suggest that there is an apparent
- 25 survival benefit in a non-randomised trial of
- 26 patients receiving radiolabelled 791T/36 antibody.
- 27 The dose of radiolabel reaching the tumour is well
- 28 below the level required to elicit tumour killing as
- 29 a result of the radiolabel alone. It is therefore

more likely that the antibody is inactivating CD55,

2	allowing complement attack of residual tumour. As
3	these patients only received a single intravenous
4	dose of 791T/36 antibody the apparent survival
5	benefit is very dramatic. Repeat injection with a
6	humanised 791T/36 antibody may have an even more
7	pronounced therapeutic benefit.
8	
9	Example 3. Production of new monoclonal antibodies
	to SCR1 and SCR2
11	
12	6-8 week old Balb/c mice were immunised twice 3
13	weeks apart by intraperitoneal injection with 791T
14	cells that over-express CD55 antigen (106 cells).
15	Mice were then boosted with SCR1-2 protein fused to
16	human Fc and purified by protein A chromatography.
17	Mice were tail bled and serum was screened for their
18	ability to recognise CD55SCR1-2/CD46SCR3-4 chimeric
19	molecules expressed by CHO cells as previously
20	described (Spendlove et al 2000 Eur J Immunol 30,
21	2944). They were also screened for their ability to
22	recognise the SCR1-2CD55Fc protein and the IC, 2N
23	and 2C peptides attached to BSA as previously
24	described (Spendlove et al 2000 Eur J Immunol 30,
25	2944). Mice producing antibodies that recognises
26	CD55SCR1 and SCR2 are boosted by an intravenous
27	injection of SCR1-2Fc protein and
28	splenocytes removed 5 days later and fused using PEG
29	with NSO myeloma cells at a 10:1 ratio. Hybridomas
30	are selected using HAT medium and screened for
31	production of antibodies recognising SRR1-2Fc
32	protein by ELISA. Hybridomas producing the correct

1 antibody are cloned by limiting dilution three times 2 a 1 cells per well to ensure clonality. The 3 monoclonal antibody is screened for its ability to recognise CD55SCR1-2/CD46SCR3-4 chimaeric molecules 4 expressed by CHO cells as previously described 5 (Spendlove et al 2000 Eur J Immunol 30, 2944). They 6 7 are also screened for their ability to recognise the SCR1-2CD55Fc protein and the IC, 2N and 2C peptides 8 attached to BSA as previously described (Spendlove 9 et al 2000 Eur J Immunol 30, 2944). To determine if 10 they recognise the same site as 791T/36 plates are 11 12 coated with CD55 as described above. They are then 13 incubated with the new monoclonal antibodies and then with biotinylated 791T/36. Binding of 791T/36 14 is quantified by avidin peroxidase and ABTS 15 substrate and the OD read at 405nm on a plate 16 17 reader. If the monoclonal antibodies recognise the same or related sites to 791T/36 they will inhibit 18 binding of 791T/36 to CD55 antigen. 19 20 All documents referred to in this specification are 21 22 herein incorporated by reference. Various 23 modifications and variations to the described embodiments of the inventions will be apparent to 24 25 those skilled in the art without departing from the scope and spirit of the invention. Although the 26 27 invention has been described in connection with specific preferred embodiments, it should be 28 understood that the invention as claimed should not 29 be unduly limited to such specific embodiments. 30 Indeed, various modifications of the described modes 31 32 of carrying out the invention which are obvious to

- those skilled in the art are intended to be covered
- 2 by the present invention.

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1 Claims

2

- 3 1. The use of (i) a naked binding member which
- 4 binds to both SCR1 and SCR2 of CD55 or (ii) a
- 5 nucleic acid encoding said binding member in the
- 6 preparation of a medicament for the neutralisation
- 7 of CD55.

8

- 9 2. The use of (i) a naked binding member which
- 10 binds to both SCR1 and SCR2 of CD55 or (ii) a
- 11 nucleic acid encoding said binding member in the
- 12 preparation of a medicament for the enhancement of
- 13 complement deposition on a tissue.

14

- 15 3. The use of (i) a naked binding member which
- 16 binds to both SCR1 and SCR2 of CD55 or (ii) a
- 17 nucleic acid encoding said binding member in the
- 18 preparation of a medicament for treating cancer.

19

- 20 4. The use according to claim 3 wherein the cancer
- 21 is one or more of colorectal, breast, ovarian,
- 22 cervical, gastric, lung, liver, skin and myeloid
- 23 (e.g. bone marrow) cancer.

24

- 25 5. The use according to any one of the preceding
- 26 claims wherein the binding member is an antibody or
- 27 a fragment thereof.

- 29 6. The use according to any one of the preceding
- 30 claims wherein the binding member binds to amino
- 31 acids 83-93and SCR2 amino acids 101-112 and amino
- 32 acids 145-157 of the sequences shown in Figure 1b.

The use according to any one of the preceding 1 7. claims wherein the binding member comprises one or 2 more of the CDRs of the antibody, or a fragment 4 thereof, produced by the cell line deposited at ATCC under accession number HB9173. 6 The use according to any one of the preceding 7 8. claims wherein the binding member is the antibody 8 791T/36 produced by the hybridoma cell deposited at 9 ATCC under accession number HB9173. 10 11. The use according to any one of claims 1 to 7 12 9. 13 wherein the binding member comprises at least one human constant region. 14 15 A naked binding member which binds to both SCR1 16 and SCR2 for use in the treatment of cancer. 17 18 A naked binding member, which binds to both 19 SCR1 and SCR2 of CD55, and an active agent as a 20 combined preparation for simultaneous, separate or 21 sequential use in the treatment of cancer. 22 23 The naked binding member according to claim 11, 24 wherein said active agent is a Doxorubicin, taxol, 25 5-Fluorouracil, Irinotecan or Cisplatin. 26 27 The naked binding member according to any one 28 of claims 10 to 12, wherein the naked binding member 29 30 is as defined in any one of claims 1 to 9.

- 1 14. A pharmaceutical composition for the treatment
- 2 of cancer, wherein the composition comprises a naked
- 3 binding member that binds to both SCR1 and SCR2 of
- 4 CD55 and a pharmaceutically acceptable excipient,
- 5 diluent or carrier.

б

- 7. 15. The pharmaceutical composition according to
- 8 claim 14, wherein the naked binding member is as
- 9 defined in any one of claims 1 to 9.

10

- 11 16. A method of neutralisation of CD55, comprising
- 12 administration of a naked binding member which
- specifically binds to SCR1 and SCR2 of CD55.

14

- 15 17. A method of enhancing complement deposition
- 16 comprising administration of a naked binding member
- which specifically binds to SCR1 and SCR2 of CD55.

18

- 19 18. A method of treating cancer comprising
- 20 administration of a therapeutically effective amount
- 21 of a naked binding member which specifically binds
- 22 to SCR1 and SCR2 of CD55 to a mammal in need
- 23 thereof.

24

- 25 19. A method according to any one of claims 16 to
- 26 18 wherein the naked binding member is as defined in
- 27 any one of claims 1 to 9.

28

- 29 20. An assay method for identification of an agent
- 30 capable of inhibiting CD55 comprising step:

. 31

a) bringing into contact a candidate agent with at 1 2 least a portion of SCR1 and SCR2 of CD55; and 3 b) 4 determining binding of said candidate agent to 5 both SCR1 and SCR2. 7 An assay method for identification of an agent 8 capable of inhibiting CD55 comprising: 9 10 (a) bringing into contact a candidate agent with at 11 least a portion of SCR1 and SCR2 of CD55 in the 12 presence of a naked binding member which in the absence of the candidate agent is capable of 13 14 binding both SCR1 and SCR2 of CD55; and 15 16 (b) determining the extent to which the candidate 17 agent inhibits binding of the naked binding member to SCR1 and SCR2 of CD55. 18 19 20 The assay method according to claim 21 wherein 21 the binding member is as defined in any one of 22 claims 6 to 9. 23 24 The assay method according to any one of claims 20 to claim 22 further comprising step (c) selecting 25 26 a candidate agent which bind both SCR1 and SCR2 of 27 CD55; and/or step (d) determining the amount of 28 complement deposition on a cell sample in the 29 presence and absence of the candidate agent. 30 31 The assay method according to any one of claims 20 to 23 wherein said portion of SCR1 and SCR2 of 32

- 1 CD55 comprises amino acids 83-93, 101-112 and 145-
- 2 157 of the sequences shown in Figure 1b.

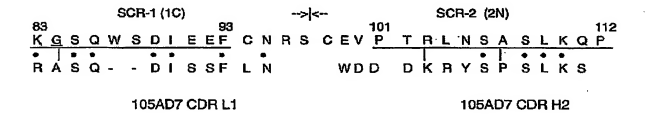
3

- 4 25. Use of an agent identified by the assay method
- of any one of claims 20 to 24 in the manufacture of
- a medicament for the treatment of cancer.

Figure 1a. CDR amino acid sequences of 105AD7 anti-idiotypic antibody

CDR	Kappa (L)	Heavy (H)
1	itcRASQDISSFLNwyq	ntSGVCVGwi
2	liyAASILQSgvp	wiaHIYWDDDKRYSPSLKSrit
3	yycQQSYKTPPSfgq	caqVLYYDFWSGYLEYFAYwgq

Figure 1b. Alignment of CDRs with CD55



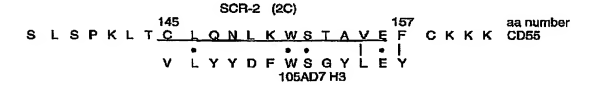
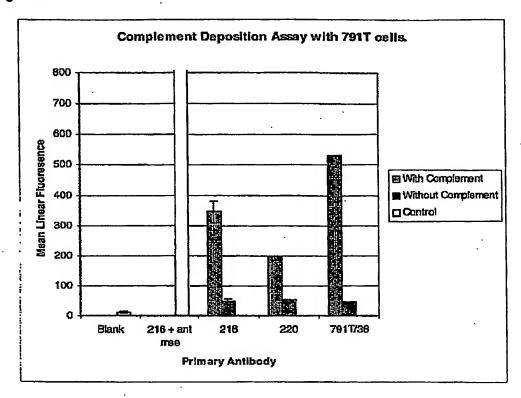


Figure 2



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